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Getting a GRiPP on everyday schistosomiasis: experience from Zimbabwe

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26 SUMMARY

27 Schistosomiasis, commonly known as bilharzia, is a parasitic disease prevalent in
28 Africa, Asia and South America. The majority of the cases occur in Sub-Saharan
29 Africa where schistosomiasis is a major public health problem impacting on child
30 health and development as well as adult health when infections become chronic.
31 Control of schistosomiasis is by treatment of infected people with the antihelminthic
32 drug praziquantel. Current schistosome control programmes advocated by the World
33 Health Assembly in 2001 are aimed at regular school based integrated deworming
34 strategies in order to reduce development of severe morbidity, promote school health
35 and to improve cognitive potential of children. Several countries in Africa have now
36 embarked on national scale deworming programmes treating millions of children
37 exposed to schistosomiasis in endemic areas without prior diagnosis of infection
38 through Mass Drug Administration (MDA) programmes. Implementing such control
39 programmes requires a concerted effort between scientists, policy makers, health
40 practitioners and several other stake holders and of course a receptive community.
41 This paper considers the contributions to global schistosome control efforts made by
42 research conducted in Zimbabwe and the historical context and developments
43 leading to the national schistosomiasis control programme in Zimbabwe giving an
44 example of Getting Research into Policy and Practice (GRiPP).

45

46

47 Key words: schistosomiasis, bilharzia, mass drug administration (MDA), Zimbabwe

48

49

50 INTRODUCTION

51 Schistosomiasis is an ancient disease, recently detected in a 5000 year old Egyptian
52 mummy (Matheson *et al* , 2014). One of its symptoms, bloody urine, is referred to in
53 a substantial number of surviving Egyptian papyri. The worm that causes urogenital
54 schistosomiasis was discovered in 1851 by Theodor Bilharz, a German physician
55 while conducting an autopsy in Egypt. He named the worm *Distomum haematobium*
56 (Foster, 1965). In 1856 Heinrich Meckel von Hemsbach proposed that the organism
57 be renamed *Bilharzia haematobium*, and the name *Schistosoma haematobium* was
58 adopted two years later in 1858. Subsequently in 1915, Leiper made the distinction
59 between *S. mansoni* and *S. haematobium* (Farley, 1991) causative agents of
60 intestinal and urogenital schistosomiasis respectively. Schistosomiasis has been
61 proposed as the explanation of the biblical curse of Jericho (Hulse, 1971), and
62 schistosome control efforts in China inspired the poem 'Farewell to the God of
63 Plague' by Mao Tse Tung (Zedong, 2007). However, schistosomiasis control in
64 present day endemic areas inspires urgent and concerted efforts to make a
65 sustained and long lasting impact on the spread and extent of the disease.

66

67 *Life cycle*

68 Schistosomes are digenetic trematodes, with two reproductive stages, one sexual
69 and the other asexual. Schistosomes have several vertebrate hosts, but only three of
70 the parasite species are important in man. Humans are the only significant definitive
71 host of *S. haematobium*, the causative agent of urogenital schistosomiasis, although
72 the infection has been found naturally in baboons and monkeys in east Africa,
73 rodents in Kenya and southern Africa, pigs in Nigeria and chimpanzees in West
74 Africa (Soulsby, 1986). Intestinal schistosomiasis is caused by *S. mansoni* and *S.*

75 *japonicum* (life cycle also described by Leiper (Leiper & Atkinson, 1915)), by adult
76 stage worms residing in the mesenteric arteries. *S. mansoni*'s most significant host is
77 the human, while *S. japonicum* is a zoonotic infection, affecting man and several
78 animals including bovines and porcines. This means that control efforts for *S.*
79 *japonicum* need to include the animal reservoirs that share habitats with man.

80 The general schistosome life cycle summarised in Soulsby (Soulsby, 1986) is
81 as follows: adult worms in copula reside in the posterior mesenteric arteries and
82 eggs are laid in the walls of the bladder, ureters and urethra (*S. haematobium*) or in
83 the intestinal mesenteric arteries (*S. mansoni* and *S. japonicum*) although *S.*
84 *haematobium* has also been demonstrated in the intestinal niches though autopsy
85 studies and excretion of the parasite's eggs in stool (Jordan *et al.*, 1993). The eggs
86 are passed out in urine or stool and will hatch in water in response to a lower
87 osmotic potential, producing miracidia. Miracidia infects the intermediate host, a
88 freshwater snail (*Bulinus globosus* for *S. haematobium*, *Oncomelania* spp. for *S.*
89 *japonicum* and *Biomphalaria* spp. for example, *Biomphalaria glabrata*), developing
90 into a mother sporocyst usually near its point of entry and after 2 weeks produces
91 daughter sporocysts. This reproduction is asexual and lasts for about 6 to 7 weeks,
92 during which daughter sporocysts migrate to other organs of the snail. Work in
93 Zimbabwe has shown that cercaria shedding is seasonal in some areas with
94 sporocysts exhibiting dormancy in winter (Shiff *et al.*, 1975). In addition, the same
95 study also reported that the pre-patent period is prolonged in winter, leading to more
96 infection in the summer months. Cercariae, the stage infective to humans, will start
97 to emerge from the daughter sporocysts 4 weeks after the initial penetration by the
98 miracidium. These are attracted to unsaturated fatty acids in the skin lipids and
99 digest their way through the skin of an exposed person, losing their tails in the

process to become schistosomulae. The schistosomulae then migrate to the lungs and eventually to the mesenteric arteries (in the case of *S. mansoni* or *S. japonicum*) or venous bladder plexus (in the case of *S. haematobium*) where they mate for life. Mated females begin to lay eggs while unmated females do not reach sexual maturity. Some of the eggs produced will leave the body through urine to repeat the life cycle. The life expectancy of adult worms is between 3 and 7 years (Fulford *et al.*, 1995).

A few features of this life cycle are worth noting: similar to other helminth macroparasites, the parasite load in the human host increases only by (re)infection (Anderson & May, 1992) through exposure to infective water, an important consideration for control programmes. The mating system is generally assumed to be monogamous but some cases of polygamy have been reported (Armstrong, 1952; Tchuem Tchuente *et al.*, 1996). It has been suggested that mating might be sequential rather than lifelong, allowing some male worms to be 'unfaithful' so that a female worm can be fertilised by more than one male in the same host (Tchuem Tchuente *et al.*, 1996). This means that as long as there is a high population of females, few males can sustain transmission. Hence, sex-differences in drug sensitivity should be a serious consideration when developing anti-schistosome drugs.

SCHISTOSOMIASIS IN ZIMBABWE

Most of the early work in African schistosomiasis was undertaken in Egypt due to the strategic importance of Egypt and the Suez Canal for imperial trade. In addition, fears raised by the incorrect hypothesis that bilharzia-causing worm were passed directly from man to man, created the political and scientific conditions that lead to

Leiper's attachment to the British Army and his important discoveries about the worms and their life cycles in Egypt (Farley, 1991). In Zimbabwe, it is quite likely that schistosomiasis was endemic before the advent of colonisation in 1890 albeit at lower prevalence at the time. In 1907, Francisco Manchego reported 'an extraordinary frequency of cases' of *S. haematobium* in the Zambezi basin, 'almost equal to that in Egypt' which had the highest infection prevalence in Africa at the time (Farley, 1991). As early as this, Manchego had observed that it was mostly children that were infected. However, it was not until Orpen described the results of a small survey in 1915 that local infections were verified (Orpen, 1915). He reported a prevalence of 31% of urinary infections among 592 African prisoners in the Salisbury jail.

Before the First World War, tropical medicine was focused mainly on the health of British colonial officials and army personnel, but, after the war, economic factors began to play an increasingly important role. Profits generated by mines in Southern Rhodesia (present day Zimbabwe) seemed threatened by workers rendered inefficient by bilharzia (Van Onselen, 1976). The Annual Public Health Reports of Southern Rhodesia showed a gradual increase in infection prevalence from 1915 to 1940 and reports following from these three and a half decades also showed an increase in infection intensity. This increase in infection prevalence and intensity appears to have been due to two major developments. First, the development in agricultural practices necessitated construction of large artificial reservoirs of water for use during the dry periods. These reservoirs provided a habitat for the intermediate hosts and allowed their populations to increase (Shiff, 1964a; Shiff, 1964b). The second development was the resettlement of the

149 autochthonous people in restricted areas such as farms and mines where the
150 populations grew without adequate sanitary systems or safe water.

151 As the prevalence of the disease was not very high in the European settlers,
152 not much work was carried out on bilharzia in the first two decades of the century
153 and the 1921 Public Health Report noted that the treatment of bilharzia in white
154 school children by tartar emetic had ‘... robbed it of much of its danger’ (Ministry of
155 Health Southern Rhodesia 1922). However, the danger of contracting the disease
156 from the Africans was ever present as the 1923 report noted: bilharzia ‘....seems
157 under control among Europeans, though of course the natives are commonly
158 infected’ (Ministry of Health Southern Rhodesia 1924). These diseased Africans
159 were a menace as they prevented the whites from enjoying their right to swim in
160 safety, and attacking the disease, the same report noted ‘might help to free the
161 country of infection and enable us to bathe safely’.

162 In 1927, William Blackie, a helminthologist from the London School of Tropical
163 Medicine and future director of the Rhodesian Public Health Laboratory arrived in
164 Rhodesia to investigate the helminth infections in the colony. A preliminary survey of
165 white children showed that bilharzia was the most serious helminth infection in the
166 country and, given the threat posed by African ‘reservoirs of disease’, a massive
167 helminth survey of the African Reserves (restricted areas where the indigenous
168 African were re-settled) was called for. His survey of the African population revealed
169 *S. haematobium* to be present with a prevalence reaching over 20% in some areas
170 (Blackie, 1932). This increase in infection was obviously evident to the health
171 authorities in Rhodesia and it prompted the establishment of a specialised laboratory
172 working on schistosomiasis in 1939.

Vic Clarke, Clive Shiff, Michael Gelfand and Dyson Blair were amongst the most prominent workers on schistosomiasis in Rhodesia. Clarke described the distribution of schistosome infection in communities, showing age-related differences in infection prevalence and intensity (Clarke, 1966) extending the work of Fisher in Zaire in the 1930s (Fisher, 1934). The studies of Clarke and Fisher suggested that the distribution of schistosome infections in human populations i.e. high infections in children which dropped in adulthood, was due to the development of protective acquired immunity; making these the first descriptions of schistosome immunology. In his studies, Clarke reported prevalences as high as 84% in some areas with 7-9 year old children having an infection prevalence as high as 98% (Clarke, 1966). Clarke went on to become the Director of the specialised laboratory where he carried out a substantial amount of work on bilharzia as well as trialling and implementing different control measures through to the late 1980's. Several researchers currently working on schistosomiasis including myself, had the privilege of being trained/taught by Clarke at the Blair Research Institute or at the University of Zimbabwe.

The specialised laboratory set up in 1939 was named the Blair Research Institute in Salisbury (now Harare) after Dr. Dyson Blair, who had been Secretary of Health in the country. A second laboratory contributing to work on schistosomiasis was the De Beers Research Laboratory established in 1965 in Chiredzi. Both laboratories form part of the research wing of the Ministry of Health and continue to work on schistosomiasis in Zimbabwe as the National Institute of Health Research under the leadership of Susan Mutambu. It is in collaboration with this institute that most of the studies described here, including our own, were conducted.

ZIMBABWE'S LEGACY TO BILHARZIA RESEARCH AND CONTROL

Research from Zimbabwe has made a significant scientific impact on our current understanding of schistosome epidemiology, the nature and development of acquired immunity, the effects of treatment on schistosome specific immune responses, the efficacy and safety of antihelminthic drugs and schistosome pathology. Workers in the 1980s and 1990s including Stephen Chandiwana, Moses Chimbari, Jerichias Ndamba, Patricia Ndhlovu and Mark Woolhouse conducted extensive studies on these several aspects of bilharzia as detailed below at the Blair Research Institute (now NIHR) and also trained several of the current generation of schistosomiasis and helminth researchers.

Studies trialling complementary control strategies including mollusciciding, biological vector control, water and sanitation (WASH) and engineering solutions have also been conducted in Zimbabwe (Chandiwana *et al.*, 1988; Chandiwana *et al.*, 1991; Chimbari, 1991; Chimbari & Ndlela, 2001; Shiff, 1970; Shiff & Kriel, 1970). These studies have had both local and global impact informing policy, design of intervention strategies as well as implementation of interventions.

Schistosome epidemiology

Schistosome epidemiology refers to the description and analysis of the patterns of transmission, infection and disease in defined populations. Understanding the epidemiology of any disease is the foundation of control strategies. It is essential to determine who is infected, where they get infection and how they transmit it. It is also essential to determine levels of infection and to understand factors influencing infectiousness and transmission. Such concepts elegantly summarised in the work of Anderson and May (Anderson & May, 1992) and Woolhouse (Woolhouse, 1998) rely

on the field studies conducted in surveys undertaken in endemic areas and hospitals. The work by Clarke for his PhD thesis (Clarke, 1966) in the then Rhodesia was central in demonstrating that children carried the heaviest infections. This, with Fisher's early work (Fisher, 1934) underlie the current WHO recommendations of targeting schistosome control programmes at primary school children (Organisation, 2002).

Epidemiological studies on the snail intermediate host conducted by Woolhouse and Chandiwana in Zimbabwe indicated the heterogeneity in human water contact behaviour which exposed them to infective water as well as heterogeneity in the snail populations at different water contact sites (Woolhouse & Chandiwana, 1989; Woolhouse & Chandiwana, 1990a; Woolhouse & Chandiwana, 1990b; Woolhouse & Chandiwana, 1990c; Woolhouse & Chandiwana, 1992; Woolhouse *et al.*, 1990). By indicating that not all people were equally exposed to infection and that not all contact points were equally infectious; these studies added to the evidence for targeted control strategies. Furthermore, habitat and ecology studies of snails in Zimbabwe by Chimbari and others, allowed for engineering interventions against schistosomiasis (Chimbari *et al.*, 1997; Chimbari *et al.*, 1996).

Schistosome immunology

The studies of Fisher and Clarke (Clarke, 1966; Fisher, 1934) suggested that protective acquired immunity developed naturally in people exposed to schistosome infection. This laid the foundation for the work by Woolhouse in 1991 in Zimbabwe (Woolhouse *et al.*, 1991) which indicated that the rate of development of this protective immunity depended on the schistosome transmission dynamics, so that in areas of high transmission, protective immunity developed quicker than in areas of

low transmission leading to infection levels peaking at higher levels and in earlier ages in the former compared to the latter, a phenomenon termed the peak shift (Woolhouse, 1992). In subsequent years while working on Zimbabwean populations, we were able to demonstrate the immunological processes underlying the peak shift and further identify immune responses associated with protection against schistosome infection (Mutapi *et al.*, 1997). We further demonstrated that antihelminthic treatment with the drug praziquantel (PZQ) accelerated the rate of development of these immune responses (Mutapi *et al.*, 1998) and we have also demonstrated that these immune responses are protective against re-infection (Bourke *et al.*, 2013). Taken together, our immuno-epidemiology studies showed that the effects of praziquantel treatment extended beyond the transient reduction of infection levels and this has been an important aspect of our recommendation for the use of PZQ in treating schistosome infection. We have also used this information in predicting the long-term effects of MDA programmes for schistosome control, highlighting the need for sustained control efforts if we are to avoid infection and disease rebounds in schistosomiasis (Mitchell *et al.*, 2014). We described the very first immunology co-infection study between urogenital schistosomiasis and *Plasmodium falciparum* malaria in our studies from Zimbabwe, and our co-infection study contributed to the increase in human schistosome -*Plasmodium* co-infection studies (Mutapi *et al.*, 2000).

Schistosomiasis the disease

In young children schistosomiasis causes abdominal pain, diarrhoea and blood in the stool (intestinal schistosomiasis), blood in urine and painful urination (urogenital schistosomiasis), nutritional deficiencies, anaemia, and decreased physical

performance and growth retardation. Clinical manifestations of chronic schistosomiasis include liver and or spleen enlargement, and the disease is frequently associated with an accumulation of fluid in the peritoneal cavity and hypertension of the abdominal blood vessels. In the case of urogenital schistosomiasis, fibrosis of the bladder and ureter, bladder cancer and kidney damage can occur. In women, urogenital schistosomiasis may present with genital lesions, vaginal bleeding, pain during sexual intercourse and nodules in the vulva and recent studies have suggested that this manifestation of schistosomiasis in females i.e. female genital schistosomiasis, may predispose to HIV infection (Christinet *et al.*, 2016). When adequately treated during childhood with praziquantel (PZQ), the antihelminthic drug of choice, these disease symptoms can be reversed (King, 2006). Early studies in Zimbabwe clearly demonstrated the classical symptoms of blood in urine. Michael Gelfand published extensively on the clinical and disease manifestations of schistosomiasis (Gelfand, 1948; Gelfand, 1963; Gelfand, 1964; Gelfand, 1966; Gelfand, 1985) and in 1950 published the influential book *Schistosomiasis in South-Central Africa* (Gelfand, 1950). Currently in the urogenital schistosomiasis field, there are calls to recognize and treat female genital schistosomiasis, (Christinet *et al.*, 2016) a condition Gelfand identified and described (Gelfand *et al.*, 1971). The next time this aspect was studied extensively was in the 2000s in Zimbabwe by a team from the then Blair Research Institute, led by Patricia Ndhlovu and collaborators from Denmark (Ndhlovu *et al.*, 2007). Gelfand went on to found the *Central African Journal of Medicine* with Joseph Ritchken in 1955. In 1962, Gelfand joined the then University of Rhodesia as the founding Professor of African Medicine. Following in this tradition of characterizing and diagnosing schistosome-related disease and morbidity, our group has been focusing on young

children and we have been particularly interested in describing the clinical manifestations of schistosomiasis in pre-school children in order to inform disease quantification and diagnosis (Wami *et al.*, 2015).

Antihelminthic treatment

The oldest recorded anti-schistosome drug is antimony potassium tartrate or tartar emetic dating back to 1605 (Duffin & Rene, 1991). From the late 1950s through the early 1980s, schistosome-infected people were treated with repeated injections of tartar emetic. In Zimbabwe, then Rhodesia, tartar emetic was used to treat school children. The impracticalities of this method of treatment were highlighted by Clarke who conducted trials of hycanthone (Etrenol Winthrop). Clarke published a Target Product Profile (TPP) for schistosome antihelminthic drug which is still applicable today (Clarke *et al.*, 1969). He indicated that a schistosome drug for mass treatment needed to be oral rather than injectable, it needed to have little or no side effects for compliance, and a single dose would be preferable to multiple doses. Praziquantel discovered in 1972 by Bayer and at the same time synthesised by Merck (Germany) fits this TTP.

Paediatric schistosomiasis

Our contribution to the antihelminthic treatment of schistosomiasis has been through the studies of the need, safety and efficacy of PZQ in preschool children. Current global initiatives from Partners of Parasite Control including the World Health Organization (WHO), Bill and Melinda Gates Foundation, UNICEF, Schistosome Control Initiative and the World Bank have been advocating regular school-based de-worming strategies in order to reduce development of severe morbidity, promote

school-child health and improve cognitive potential of children. Praziquantel is being used for treating children in Africa through several governmental and non-government initiatives for example, the Schistosome Control Initiative. Until recently, schistosomiasis in preschool children was a largely ignored problem in terms of control as a result of several reasons including (a) a lack of data on their exposure to infection, (b) unknown levels of infection and morbidity in this age group, (c) unknown safety in this age group (the original safety studies in the 1970s were conducted in children aged 5 years and above and (d) unknown efficacy of the drug in this age group (Mutapi *et al.*, 2011; Stothard & Gabrielli, 2007; Stothard *et al.*, 2013). Through a series of studies in Zimbabwe (Mutapi *et al.*, 2011) we joined a group of scientists who conducted studies in pre-school children to collect the evidence base to refute the four points raised above (World Health Organisation 2012). The work culminated in 2012 in changes in WHO guidelines for the treatment of paediatric schistosomes (World Health Organisation 2012). We and others also called for a child-appropriate formulation of PZQ, an appeal that was taken up by the private public partnership named the Paediatric Praziquantel Consortium (World Health Organisation 2012). It is indeed encouraging to see the recent announcement from this Consortium (<http://www.pediatricpraziquantelconsortium.org/news-events/news.html>) that a potential paediatric praziquantel tablet has commenced phase II clinical trials in the Ivory Coast.

WASH strategies

As is clear from the life cycle of schistosomiasis, fresh water plays a critical role for the maintenance of the life cycle and transmission to humans- upon reaching fresh water, eggs from human urine or stool hatch into the stage infective to the

348 snail intermediate hosts. Hence, poor sanitation allows the contamination of
349 water sources with the parasites. People become infected when they come into
350 contact with fresh water where the snails have shed the infective cercariae. This
351 usually happens during swimming, bathing or collection of water for domestic
352 use in rivers. Hence, provision of safe water for domestic use would reduce
353 transmission of the parasites to humans. Unfortunately, the global distribution of
354 schistosomiasis overlaps with the areas where some of the poorest populations
355 inhabit. This means that safe water and sanitation provisions are poorest in
356 these areas. The challenge is to provide appropriate (water, sanitation and
357 hygiene) WASH technologies (Steinmann *et al.*, 2006). Zimbabwe has been at
358 the forefront of developing and implementing such technologies. The Blair Toilet
359 (named after the Blair Research Institute where it was developed) or Ventilation
360 Improved Pit (VIP) latrine developed by Peter Morgan in the 1970s is an
361 outstanding example of these efforts. This is a toilet built with local materials and
362 based on the design of turrets which allows airflow into the toilet, but stops
363 smells and flies escaping (see Figure 1). Peter Morgan also popularised the the
364 Bush Pump, a reliable simple lever action water pump made using local
365 components that can be operated by all age groups to get water from a
366 protected well (see Figure 2 showing children using a borehole constructed from
367 the bush pump design) which was originally designed by the water engineer
368 Tommy Murgatroyd in the Southern Rhodesia Ministry of Water Development in
369 1933 (de Laet and Mol, 2000). The pit latrine and bush pump have been adopted
370 in Zimbabwe and elsewhere across Africa and as recognition of the global
371 impact this toilet and pump designs, Peter Morgan received the Stockholm
372 Water Prize in 2013. Nonetheless, even with these appropriate local

technologies, there is still a large population of rural Zimbabweans not utilising the available pit latrines or building boreholes. The issue then is not about access or availability but rather, human behaviour and social context. There is need to understand the drivers of this human behaviour and come up with solutions to 'nudge' people towards the use of toilets and safe water sources.

Snail control

Various intervention studies have been trialled in Zimbabwe with differing levels of success (Chimbari, 1991; Chimbari *et al.*, 1992; Chimbari & Ndlela, 2001). Integrated snail vector control and antihelminthic treatment of infected people was implemented in Kariba in 1967 after the filling of Kariba Dam, the source of hydroelectric power for Zimbabwe and Zambia (Chimbari, 2012). Snail control was the strategy recommended by the World Health Organisation at the time and was achieved by application of the molluscicide niclosamide and habitat destruction (removal of the water weed *Salvinia sp.*) with antihelminthic treatment targeted at infected people. In the 1950s copper sulphate and sodium pentachlorophenate were the chemical molluscicides used, they were replaced by niclosamide believed to be less toxic to humans, cattle, plants and other aquatic life (Foster *et al.*, 1960). In the 1960s and 1970s, several mollusciding studies were conducted in Zimbabwe. In the case of work by Shiff and colleagues, the earlier studies showing seasonality in the transmission of schistosomiasis in some regions of the country (Shiff *et al.*, 1975) meant that this knowledge could be utilised in designing mollusciding approaches. Thus, they showed that *S. haematobium* transmission could be significantly reduced by annual mollusciding with Bayluscide® (a formulation of niclosamide) in winter to kill off the intermediate host snails (Shiff *et al.*, 1979). Shiff and colleagues including

Clarke also demonstrated that mollusciding of irrigation canals using niclosamide drip-feed methods every 6-8 months as well as regular treatment of drains with the molluscicide also significantly reduced the risk of infection with *S. mansoni* and *S. haematobium* in children (Shiff *et al.*, 1973). They also conducted an economic costing of the work as well as operational assessment concluding that only 10% level of surveillance and incidence check of sentinel sites within the irrigation was sufficient to provide informative monitoring and evaluation of the efficacy and long-term effects of the molluscicide control efforts (Shiff *et al.*, 1973). In a recent meta-analysis of 35 molluscicide studies including several from Zimbabwe, King and colleagues reported that on average mollusciding reduced the odds of infection by 77% with the effects increased if mollusciding was integrated with antihelminthic treatment of the human population, while incidence was reduced by 64%, but interestingly antihelminthic treatment did not influence the incidence of infection (King *et al.*, 2015).

The irrigation mollusciding study in Zimbabwe in 1973 showed that controlling schistosomiasis via this method cost between USD54,800 and USD 55,500 for a 30,000 ha irrigation scheme (Shiff *et al.*, 1973). Thus, although the control efforts integrating antihelminthic treatment and mollusciding proved effective, not surprisingly, the vector control was not sustainable, either economically or environmentally. To overcome the toxic effects of niclosamide on plants, other snails (potential competitors) and fish, biological control strategies presented an attractive alternative. In Zimbabwe different biological interventions have been investigated; these included molluscicides derived from the plants *Phytolacca dodecandra* and *Jatropha curcas* with mixed efficacy and mixed community uptake (Madhina *et al.*,

1996). In addition snail predators have also been invoked in the form of ducks and fish (Chimbari, 2012). Poaching of the former (which were non-indigenous duck species) and low efficacy of the latter reduced the uptake of these interventions. Introduction of non-host competitor snails did not have a significant effect on the population of the intermediate host snails (Chimbari, 2012).

Engineering strategies to control schistosomiasis

Engineering and environmental interventions cannot be stressed enough in schistosome control. Long before the schistosome epidemic in Senegal in the 1990s, following the damming of the Senegal river to provide water for a sugar irrigation scheme in Richard Toll (Stelma *et al.*, 1993), Zimbabwe had experienced a schistosome epidemic as a result of the construction of the Kariba Dam in the late 1950s (Hira, 1970). The lessons learnt from this episode inspired the collaborative work between health professionals and engineers in the design of dam and irrigation schemes. A very successful example of this was the Mushandike Irrigation Scheme initiated in 1986 (Chimbari, 1991). Irrigation canals were lined to facilitate fast movement of irrigation water to dislodge snails and also comprised features which flushed out and trapped snails. Toilets were constructed along the scheme in a matrix ensuring that the workers were always nearer to a toilet than to the bush (Chimbari, 2012). This programme was so successful that despite the high costs of the design, the model was adopted by the Department of Irrigation in Zimbabwe as standard for all small irrigation schemes (Chimbari, 2012).

ZIMBWBWE'S NATIONAL SCHISTOSOMIASIS CONTROL PROGRAMME

447 With the history of commitment to schistosome research and control demonstrated
448 by the previous studies, it is not surprising that Zimbabwe has conducted regular
449 national surveys of schistosome infection in humans as well as the distribution and
450 infectivity of intermediate host snails. The first comprehensive national
451 schistosomiasis survey was conducted in 1982, followed by the second in 1992 (see
452 (Chimbari, 2012) for details). We conducted the third and most recent national
453 schistosomiasis and soil transmitted helminth (SHT) survey in Zimbabwe in 2010
454 (Midzi *et al.*, 2014). One of the changes that occurred in the intervening period
455 between the 1992 survey and our survey was the momentum from the global
456 initiative to control schistosomiasis following the 2001 World Health Assembly
457 Resolution 54.19 to treat at least 75% of all school aged children who are at risk of
458 morbidity from schistosomiasis and Soil transmitted helminths (STH) by the year
459 2010. This, coupled with the wider availability and significant reduction in price of the
460 antihelminthic drug PZQ, made for a suitable environment to implement a national
461 schistosomiasis control programme in Zimbabwe. When we published the findings of
462 our national schistosomiasis survey, we also included a national plan of action
463 incorporating the treatment strategies recommended by the WHO (Midzi *et al.*,
464 2014). Following the national helminth survey, we contributed to the formulation of
465 the national schistosomiasis and helminth control policy in Zimbabwe through a
466 workshop hosted by the Ministry of Health in Zimbabwe. To continue with the
467 research legacy of Zimbabwe and inter-sectorial collaborative approaches to
468 schistosome control, this policy document highlighted among others, the importance
469 of continued scientific research, dialogue between engineers and health
470 professionals, dialogue between the ministries of health and education both for
471 health education and implementation of the national control programmes and

community involvement. Following operational results from studies we conducted in schools in Zimbabwe and input from several stakeholders the Ministry of Health in Zimbabwe launched a the 5-year national schistosomiasis and soil transmitted control programme in September 2012. This programme is targeting just under 5 million primary school children throughout the country, treating all children annually regardless of the transmission level in the areas. The full evaluation of the programme will be conducted at the end of the 5 year programme but early indications from the sentinel monitoring and evaluation sites are that there has been a reduction in infection levels.

FUTURE PLANS AND CONCLUSION

The question for Zimbabwe and all other countries currently implementing national schistosomiasis control programmes is what happens after the 5 years of MDA? From our immuno-epidemiology and quantitative studies in Zimbabwe, we have predicted that cessation of MDA programmes may result in a rebound in infection to levels higher than pre-treatment levels (Mitchell *et al.*, 2014). Indeed earlier studies in Zimbabwe in the 1990s showed infection levels returning to pre-intervention levels when control (even strategies using integrated methods applied for 5-year periods were ceased) (Chimbari, 2012). These studies indicate that there is need for sustained control efforts and long-term planning to avoid areas of refugia for the parasites as well to facilitate the move from morbidity control to controlling transmission. There is also need for inclusive controls strategies if elimination is a realistic goal for schistosomiasis. Targeting primary school children while leaving the preschool children and adults will not lead to elimination of the diseases especially

where untreated infections in adulthood can become chronic with complications only addressed by surgery.

There are still areas needing more research including diagnostics, therapeutics and operational aspects as recently highlighted by Secor (Secor & Montgomery, 2015). There is also need for a more integrated approach to disease control involving dialogue between different sectors such as social scientists, engineers, architects (to enable building of human cities and dwellings that interrupt parasite transmission) and economists to come up with sustainable solutions to schistosome control. The current generation of schistosome researchers working in Zimbabwe aims to contribute to this knowledge base and strengthen the legacy of putting research before policy and implementation in schistosomiasis control.

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523

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527

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For Peer Review

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752 LIST OF FIGURES

753

754 Fig 1: Photographs of VIP Latrines from Zimbabwe A) family setting, B) school
755 setting

756

757 Fig 2: Photograph of a borehole using the bust pump design (village setting)

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For Peer Review

Getting a GRiPP on everyday schistosomiasis: experience from Zimbabwe

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Running title: History of Schistosomiasis Research in Zimbabwe

26 SUMMARY

27 Schistosomiasis, commonly known as bilharzia, is a parasitic disease prevalent in
28 Africa, Asia and South America. The majority of the cases occur in Sub-Saharan
29 Africa where schistosomiasis is a major public health problem impacting on child
30 health and development as well as adult health when infections become chronic.
31 Control of schistosomiasis is by treatment of infected people with the antihelminthic
32 drug praziquantel. Current schistosome control programmes advocated by the World
33 Health Assembly in 2001 are aimed at regular school based integrated deworming
34 strategies in order to reduce development of severe morbidity, promote school health
35 and to improve cognitive potential of children. Several countries in Africa have now
36 embarked on national scale deworming programmes treating millions of children
37 exposed to schistosomiasis in endemic areas without prior diagnosis of infection
38 through Mass Drug Administration (MDA) programmes. Implementing such control
39 programmes requires a concerted effort between scientists, policy makers, health
40 practitioners and several other stake holders and of course a receptive community.
41 This paper considers the contributions to global schistosome control efforts made by
42 research conducted in Zimbabwe and the historical context and developments
43 leading to the national schistosomiasis control programme in Zimbabwe giving an
44 example of Getting Research into Policy and Practice (GRiPP).

45

46

47 Key words: schistosomiasis, bilharzia, mass drug administration (MDA), Zimbabwe

48

49

50 INTRODUCTION

51 Schistosomiasis is an ancient disease, recently detected in a 5000 year old Egyptian
52 mummy (Matheson *et al* , 2014). One of its symptoms, bloody urine, is referred to in
53 a substantial number of surviving Egyptian papyri. The worm that causes urogenital
54 schistosomiasis was discovered in 1851 by Theodor Bilharz, a German physician
55 while conducting an autopsy in Egypt. He named the worm *Distomum haematobium*
56 (Foster, 1965). In 1856 Heinrich Meckel von Hemsbach proposed that the organism
57 be renamed *Bilharzia haematobium*, and the name *Schistosoma haematobium* was
58 adopted two years later in 1858. Subsequently in 1915, Leiper made the distinction
59 between *S. mansoni* and *S. haematobium* (Farley, 1991) causative agents of
60 intestinal and urogenital schistosomiasis respectively. Schistosomiasis has been
61 proposed as the explanation of the biblical curse of Jericho (Hulse, 1971), and
62 schistosome control efforts in China inspired the poem 'Farewell to the God of
63 Plague' by Mao Tse Tung (Zedong, 2007). However, schistosomiasis control in
64 present day endemic areas inspires urgent and concerted efforts to make a
65 sustained and long lasting impact on the spread and extent of the disease.

66

67 *Life cycle*

68 Schistosomes are digenetic trematodes, with two reproductive stages, one sexual
69 and the other asexual. Schistosomes have several vertebrate hosts, but only three of
70 the parasite species are important in man. Humans are the only significant definitive
71 host of *S. haematobium*, the causative agent of urogenital schistosomiasis, although
72 the infection has been found naturally in baboons and monkeys in east Africa,
73 rodents in Kenya and southern Africa, pigs in Nigeria and chimpanzees in West
74 Africa (Soulsby, 1986). Intestinal schistosomiasis is caused by *S. mansoni* and *S.*

75 *japonicum* (life cycle also described by Leiper (Leiper & Atkinson, 1915)), by adult
76 stage worms residing in the mesenteric arteries. *S. mansoni*'s most significant host is
77 the human, while *S. japonicum* is a zoonotic infection, affecting man and several
78 animals including bovines and porcines. This means that control efforts for *S.*
79 *japonicum* need to include the animal reservoirs that share habitats with man.

80 The general schistosome life cycle summarised in Soulsby (Soulsby, 1986) is
81 as follows: adult worms in copula reside in the posterior mesenteric arteries and
82 eggs are laid in the walls of the bladder, ureters and urethra (*S. haematobium*) or in
83 the intestinal mesenteric arteries (*S. mansoni* and *S. japonicum*) although *S.*
84 *haematobium* has also been demonstrated in the intestinal niches though autopsy
85 studies and excretion of the parasite's eggs in stool (Jordan *et al.*, 1993). The eggs
86 are passed out in urine or stool and will hatch in water in response to a lower
87 osmotic potential, producing miracidia. Miracidia infect the intermediate host, a
88 freshwater snail (*Bulinus globosus* for *S. haematobium*, *Oncomelania* spp. for *S.*
89 *japonicum* and *Biomphalaria* spp. for example, *Biomphalaria glabrata*), developing
90 into a mother sporocyst usually near its point of entry and after 2 weeks produces
91 daughter sporocysts. This reproduction is asexual and lasts for about 6 to 7 weeks,
92 during which daughter sporocysts migrate to other organs of the snail. Work in
93 Zimbabwe has shown that cercaria shedding is seasonal in some areas with
94 sporocysts exhibiting dormancy in winter (Shiff *et al.*, 1975). In addition, the same
95 study also reported that the pre-patent period is prolonged in winter, leading to more
96 infection in the summer months. Cercaria, the stage infective to humans, will start to
97 emerge from the daughter sporocysts 4 weeks after the initial penetration by the
98 miracidium. These are attracted to unsaturated fatty acids in the skin lipids and
99 digest their way through the skin of an exposed person, losing their tails in the

process to become schistosomulae. The schistosomulae then migrate to the lungs and eventually to the mesenteric arteries (in the case of *S. mansoni* or *S. japonicum*) or venous bladder plexus (in the case of *S. haematobium*) where they mate for life. Mated females begin to lay eggs while unmated females do not reach sexual maturity. Some of the eggs produced will leave the body through urine to repeat the life cycle. The life expectancy of adult worms is between 3 and 7 years (Fulford *et al.*, 1995).

A few features of this life cycle are worth noting: similar to other helminth macroparasites, the parasite load in the human host increases only by (re)infection (Anderson & May, 1992) through exposure to infective water, an important consideration for control programmes. The mating system is generally assumed to be monogamous but some cases of polygamy have been reported (Armstrong, 1952; Tchuem Tchuente *et al.*, 1996). It has been suggested that mating might be sequential rather than lifelong, allowing some male worms to be 'unfaithful' so that a female worm can be fertilised by more than one male in the same host (Tchuem Tchuente *et al.*, 1996). This means that as long as there is a high population of females, few males can sustain transmission. Hence, sex-differences in drug sensitivity should be a serious consideration when developing anti-schistosome drugs.

SCHISTOSOMIASIS IN ZIMBABWE

Most of the early work in African schistosomiasis was undertaken in Egypt due to the strategic importance of Egypt and the Suez Canal for imperial trade. In addition, fears raised by the incorrect hypothesis that bilharzia-causing worm were passed directly from man to man, created the political and scientific conditions that led to

125 Leiper' s attachment to the British Army and his important discoveries about the
126 worms and their life cycles in Egypt (Farley, 1991). In Zimbabwe, it is quite likely that
127 schistosomiasis was endemic before the advent of colonisation in 1890 albeit at
128 lower prevalence at the time. In 1907, Francisco Manchego reported 'an
129 extraordinary frequency of cases' of *S. haematobium* in the Zambezi basin, 'almost
130 equal to that in Egypt' which had the highest infection prevalence in Africa at the time
131 (Farley, 1991). As early as this, Manchego had observed that it was mostly children
132 that were infected. However, it was not until Orpen described the results of a small
133 survey in 1915 that local infections were verified (Orpen, 1915). He reported a
134 prevalence of 31% of urinary infections among 592 African prisoners in the Salisbury
135 jail.

136 Before the First World War, tropical medicine was focused mainly on the
137 health of British colonial officials and army personnel, but, after the war, economic
138 factors began to play an increasingly important role. Profits generated by mines in
139 Southern Rhodesia (present day Zimbabwe) seemed threatened by workers
140 rendered inefficient by bilharzia (Van Onselen, 1976). The Annual Public Health
141 Reports of Southern Rhodesia showed a gradual increase in infection prevalence
142 from 1915 to 1940 and reports following from these three and a half decades also
143 showed an increase in infection intensity. This increase in infection prevalence and
144 intensity appears to have been due to two major developments. First, the
145 development in agricultural practices necessitated construction of large artificial
146 reservoirs of water for use during the dry periods. These reservoirs provided a
147 habitat for the intermediate hosts and allowed their populations to increase (Shiff,
148 1964a; Shiff, 1964b). The second development was the resettlement of the

149 autochthonous people in restricted areas such as farms and mines where the
150 populations grew without adequate sanitary systems or safe water.

151 As the prevalence of the disease was not very high in the European settlers,
152 not much work was carried out on bilharzia in the first two decades of the century
153 and the 1921 Public Health Report noted that the treatment of bilharzia in white
154 school children by tartar emetic had ‘... robbed it of much of its danger’ (Ministry of
155 Health Southern Rhodesia 1922). However, the danger of contracting the disease
156 from the Africans was ever present as the 1923 report noted: bilharzia ‘....seems
157 under control among Europeans, though of course the natives are commonly
158 infected’ (Ministry of Health Southern Rhodesia 1924). These diseased ‘..Africans
159 were a menace as they prevented the whites from enjoying their right to swim in
160 safety’, and attacking the disease, the same report noted ‘might help to free the
161 country of infection and enable us to bathe safely’.

162 In 1927, William Blackie, a helminthologist from the London School of Tropical
163 Medicine and future director of the Rhodesian Public Health Laboratory arrived in
164 Rhodesia to investigate the helminth infections in the colony. A preliminary survey of
165 white children showed that bilharzia was the most serious helminth infection in the
166 country and, given the threat posed by African ‘reservoirs of disease’, a massive
167 helminth survey of the African Reserves (restricted areas where the indigenous
168 African were re-settled) was called for. His survey of the African population revealed
169 *S. haematobium* to be present with a prevalence reaching over 20% in some areas
170 (Blackie, 1932). This increase in infection was obviously evident to the health
171 authorities in Rhodesia and it prompted the establishment of a specialised laboratory
172 working on schistosomiasis in 1939.

Vic Clarke, Clive Shiff, Michael Gelfand and Dyson Blair were amongst the most prominent workers on schistosomiasis in Rhodesia. Clarke described the distribution of schistosome infection in communities, showing age-related differences in infection prevalence and intensity (Clarke, 1966) extending the work of Fisher in Zaire in the 1930s (Fisher, 1934). The studies of Clarke and Fisher suggested that the distribution of schistosome infections in human populations i.e. high infections in children which dropped in adulthood, was due to the development of protective acquired immunity; making these the first descriptions of schistosome immunology. In his studies, Clarke reported prevalences as high as 84% in some areas with 7-9 year old children having an infection prevalence as high as 98% (Clarke, 1966). Clarke went on to become the Director of the specialised laboratory where he carried out a substantial amount of work on bilharzia as well as trialling and implementing different control measures through to the late 1980's. Several researchers currently working on schistosomiasis including myself, had the privilege of being trained/taught by Clarke at the Blair Research Institute or at the University of Zimbabwe.

The specialised laboratory set up in 1939 was named the Blair Research Institute in Salisbury (now Harare) after Dr. Dyson Blair, who had been Secretary of Health in the country. A second laboratory contributing to work on schistosomiasis was the De Beers Research Laboratory established in 1965 in Chiredzi. Both laboratories form part of the research wing of the Ministry of Health and continue to work on schistosomiasis in Zimbabwe as the National Institute of Health Research under the leadership of Susan Mutambu. It is in collaboration with this institute that most of the studies described here, including our own, were conducted.

ZIMBABWE'S LEGACY TO BILHARZIA RESEARCH AND CONTROL

Research from Zimbabwe has made a significant scientific impact on our current understanding of schistosome epidemiology, the nature and development of acquired immunity, the effects of treatment on schistosome specific immune responses, the efficacy and safety of antihelminthic drugs and schistosome pathology. Workers in the 1980s and 1990s including Stephen Chandiwana, Moses Chimbari, Jerichias Ndamba, Patricia Ndhlovu and Mark Woolhouse conducted extensive studies on these several aspects of bilharzia as detailed below at the Blair Research Institute (now NIHR) and also trained several of the current generation of schistosomiasis and helminth researchers.

Studies trialling complementary control strategies including mollusciciding, biological vector control, water and sanitation (WASH) and engineering solutions have also been conducted in Zimbabwe (Chandiwana *et al.*, 1988; Chandiwana *et al.*, 1991; Chimbari, 1991; Chimbari & Ndlela, 2001; Shiff, 1970; Shiff & Kriel, 1970). These studies have had both local and global impact informing policy, design of intervention strategies as well as implementation of interventions.

Schistosome epidemiology

Schistosome epidemiology refers to the description and analysis of the patterns of transmission, infection and disease in defined populations. Understanding the epidemiology of any disease is the foundation of control strategies. It is essential to determine who is infected, where they get infection and how they transmit it. It is also essential to determine levels of infection and to understand factors influencing infectiousness and transmission. Such concepts elegantly summarised in the work of Anderson and May (Anderson & May, 1992) and Woolhouse (Woolhouse, 1998) rely

on the field studies conducted in surveys undertaken in endemic areas and hospitals. The work by Clarke for his PhD thesis (Clarke, 1966) in the then Rhodesia was central in demonstrating that children carried the heaviest infections. This, with Fisher's early work (Fisher, 1934) underlies the current WHO recommendations of targeting schistosome control programmes at primary school children (Organisation, 2002).

Epidemiological studies on the snail intermediate host conducted by Woolhouse and Chandiwana in Zimbabwe indicated the heterogeneity in human water contact behaviour which exposed them to infective water as well as heterogeneity in the snail populations at different water contact sites (Woolhouse & Chandiwana, 1989; Woolhouse & Chandiwana, 1990a; Woolhouse & Chandiwana, 1990b; Woolhouse & Chandiwana, 1990c; Woolhouse & Chandiwana, 1992; Woolhouse *et al.*, 1990). By indicating that not all people were equally exposed to infection and that not all contact points were equally infectious; these studies added to the evidence for targeted control strategies. Furthermore, habitat and ecology studies of snails in Zimbabwe by Chimbari and others, allowed for engineering interventions against schistosomiasis (Chimbari *et al.*, 1997; Chimbari *et al.*, 1996).

Schistosome immunology

The studies of Fisher and Clarke (Clarke, 1966; Fisher, 1934) suggested that protective acquired immunity developed naturally in people exposed to schistosome infection. This laid the foundation for the work by Woolhouse in 1991 in Zimbabwe (Woolhouse *et al.*, 1991) which indicated that the rate of development of this protective immunity depended on the schistosome transmission dynamics, so that in areas of high transmission, protective immunity developed quicker than in areas of

low transmission leading to infection levels peaking at higher levels and in earlier ages in the former compared to the latter, a phenomenon termed the peak shift (Woolhouse, 1992). In subsequent years while working on Zimbabwean populations, we were able to demonstrate the immunological processes underlying the peak shift and further identify immune responses associated with protection against schistosome infection (Mutapi *et al.*, 1997). We further demonstrated that antihelminthic treatment with the drug praziquantel (PZQ) accelerated the rate of development of these immune responses (Mutapi *et al.*, 1998) and we have also demonstrated that these immune responses are protective against re-infection (Bourke *et al.*, 2013). Taken together, our immuno-epidemiology studies showed that the effects of praziquantel treatment extended beyond the transient reduction of infection levels and this has been an important aspect of our recommendation for the use of PZQ in treating schistosome infection. We have also used this information in predicting the long-term effects of MDA programmes for schistosome control, highlighting the need for sustained control efforts if we are to avoid infection and disease rebounds in schistosomiasis (Mitchell *et al.*, 2014). We described the very first immunology co-infection study between urogenital schistosomiasis and *Plasmodium falciparum* malaria in our studies from Zimbabwe, and our co-infection study contributed to the increase in human schistosome -*Plasmodium* co-infection studies (Mutapi *et al.*, 2000).

Schistosomiasis the disease

In young children schistosomiasis causes abdominal pain, diarrhoea and blood in the stool (intestinal schistosomiasis), blood in urine and painful urination (urogenital schistosomiasis), nutritional deficiencies, anaemia, and decreased physical

performance and growth retardation. Clinical manifestations of chronic schistosomiasis include liver and or spleen enlargement, and the disease is frequently associated with an accumulation of fluid in the peritoneal cavity and hypertension of the abdominal blood vessels. In the case of urogenital schistosomiasis, fibrosis of the bladder and ureter, bladder cancer and kidney damage can occur. In women, urogenital schistosomiasis may present with genital lesions, vaginal bleeding, pain during sexual intercourse and nodules in the vulva and recent studies have suggested that this manifestation of schistosomiasis in females i.e. female genital schistosomiasis, may predispose to HIV infection (Christinet *et al.*, 2016). When adequately treated during childhood with praziquantel (PZQ), the antihelminthic drug of choice, these disease symptoms can be reversed (King, 2006). Early studies in Zimbabwe clearly demonstrated the classical symptoms of blood in urine. Michael Gelfand published extensively on the clinical and disease manifestations of schistosomiasis (Gelfand, 1948; Gelfand, 1963; Gelfand, 1964; Gelfand, 1966; Gelfand, 1985) and in 1950 published the influential book *Schistosomiasis in South-Central Africa* (Gelfand, 1950). Currently in the urogenital schistosomiasis field, there are calls to recognize and treat female genital schistosomiasis, (Christinet *et al.*, 2016) a condition Gelfand identified and described (Gelfand *et al.*, 1971). The next time this aspect was studied extensively was in the 2000s in Zimbabwe by a team from the then Blair Research Institute, led by Patricia Ndhlovu and collaborators from Denmark (Ndhlovu *et al.*, 2007). Gelfand went on to found the *Central African Journal of Medicine* with Joseph Ritchken in 1955. In 1962, Gelfand joined the then University of Rhodesia as the founding Professor of African Medicine. Following in this tradition of characterizing and diagnosing schistosome-related disease and morbidity, our group has been focusing on young

children and we have been particularly interested in describing the clinical manifestations of schistosomiasis in pre-school children in order to inform disease quantification and diagnosis (Wami *et al.*, 2015).

Antihelminthic treatment

The oldest recorded anti-schistosome drug is antimony potassium tartrate or tartar emetic dating back to 1605 (Duffin & Rene, 1991). From the late 1950s through the early 1980s, schistosome-infected people were treated with repeated injections of tartar emetic. In Zimbabwe, then Rhodesia, tartar emetic was used to treat school children. The impracticalities of this method of treatment were highlighted by Clarke who conducted trials of hycanthone (Etrenol Winthrop). Clarke published a Target Product Profile (TPP) for schistosome antihelminthic drug which is still applicable today (Clarke *et al.*, 1969). He indicated that a schistosome drug for mass treatment needed to be oral rather than injectable, it needed to have little or no side effects for compliance, and a single dose would be preferable to multiple doses. Praziquantel discovered in 1972 by Bayer and at the same time synthesised by Merck (Germany) fits this TTP.

Paediatric schistosomiasis

Our contribution to the antihelminthic treatment of schistosomiasis has been through the studies of the need, safety and efficacy of PZQ in preschool children. Current global initiatives from Partners of Parasite Control including the World Health Organization (WHO), Bill and Melinda Gates Foundation, UNICEF, Schistosome Control Initiative and the World Bank have been advocating regular school-based de-worming strategies in order to reduce development of severe morbidity, promote

school-child health and improve cognitive potential of children. Praziquantel is being used for treating children in Africa through several governmental and non-government initiatives for example, the Schistosome Control Initiative. Until recently, schistosomiasis in preschool children was a largely ignored problem in terms of control as a result of several reasons including (a) a lack of data on their exposure to infection, (b) unknown levels of infection and morbidity in this age group, (c) unknown safety in this age group (the original safety studies in the 1970s were conducted in children aged 5 years and above and (d) unknown efficacy of the drug in this age group (Mutapi *et al.*, 2011; Stothard & Gabrielli, 2007; Stothard *et al.*, 2013). Through a series of studies in Zimbabwe (Mutapi *et al.*, 2011) we joined a group of scientists who conducted studies in pre-school children to collect the evidence base to refute the four points raised above (World Health Organisation 2012). The work culminated in 2012 in changes in WHO guidelines for the treatment of paediatric schistosomes (World Health Organisation 2012). We and others also called for a child-appropriate formulation of PZQ, an appeal that was taken up by the private public partnership named the Paediatric Praziquantel Consortium (World Health Organisation 2012). It is indeed encouraging to see the recent announcement from this Consortium (<http://www.pediatricpraziquantelconsortium.org/news-events/news.html>) that a potential paediatric praziquantel tablet has commenced phase II clinical trials in the Ivory Coast.

WASH strategies

As is clear from the life cycle of schistosomiasis, fresh water plays a critical role for the maintenance of the life cycle and transmission to humans- upon reaching fresh water, eggs from human urine or stool hatch into the stage infective to the

348 snail intermediate hosts. Hence, poor sanitation allows the contamination of
349 water sources with the parasites. People become infected when they come into
350 contact with fresh water where the snails have shed the infective cercariae. This
351 usually happens during swimming, bathing or collection of water for domestic
352 use in rivers. Hence, provision of safe water for domestic use would reduce
353 transmission of the parasites to humans. Unfortunately, the global distribution of
354 schistosomiasis overlaps with the areas where some of the poorest populations
355 inhabit. This means that safe water and sanitation provisions are poorest in
356 these areas. The challenge is to provide appropriate (water, sanitation and
357 hygiene) WASH technologies (Steinmann *et al.*, 2006). Zimbabwe has been at
358 the forefront of developing and implementing such technologies. The Blair Toilet
359 (named after the Blair Research Institute where it was developed) or Ventilation
360 Improved Pit (VIP) latrine developed by Peter Morgan in the 1970s is an
361 outstanding example of these efforts. This is a toilet built with local materials and
362 based on the design of turrets which allows airflow into the toilet, but stops
363 smells and flies escaping (see Figure 1). Peter Morgan also popularised the the
364 Bush Pump, a reliable simple lever action water pump made using local
365 components that can be operated by all age groups to get water from a
366 protected well (see Figure 2 showing children using a borehole constructed from
367 the bush pump design) which was originally designed by the water engineer
368 Tommy Murgatroyd in the Southern Rhodesia Ministry of Water Development in
369 1933 (de Laet and Mol, 2000). The pit latrine and bush pump have been adopted
370 in Zimbabwe and elsewhere across Africa and as recognition of the global
371 impact this toilet and pump designs, Peter Morgan received the Stockholm
372 Water Prize in 2013. Nonetheless, even with these appropriate local

technologies, there is still a large population of rural Zimbabweans not utilising the available pit latrines or building boreholes. The issue then is not about access or availability but rather, human behaviour and social context. There is need to understand the drivers of this human behaviour and come up with solutions to 'nudge' people towards the use of toilets and safe water sources.

Snail control

Various intervention studies have been trialled in Zimbabwe with differing levels of success (Chimbari, 1991; Chimbari *et al.*, 1992; Chimbari & Ndlela, 2001). Integrated snail vector control and antihelminthic treatment of infected people was implemented in Kariba in 1967 after the filling of Kariba Dam, the source of hydroelectric power for Zimbabwe and Zambia (Chimbari, 2012). Snail control was the strategy recommended by the World Health Organisation at the time and was achieved by application of the molluscicide niclosamide and habitat destruction (removal of the water weed *Salvinia sp.*) with antihelminthic treatment targeted at infected people. In the 1950s copper sulphate and sodium pentachlorophenate were the chemical molluscicides used, they were replaced by niclosamide believed to be less toxic to humans, cattle, plants and other aquatic life (Foster *et al.*, 1960). In the 1960s and 1970s, several mollusciding studies were conducted in Zimbabwe. In the case of work by Shiff and colleagues, the earlier studies showing seasonality in the transmission of schistosomiasis in some regions of the country (Shiff *et al.*, 1975) meant that this knowledge could be utilised in designing mollusciding approaches. Thus, they showed that *S. haematobium* transmission could be significantly reduced by annual mollusciding with Bayluscide® (a formulation of niclosamide) in winter to kill off the intermediate host snails (Shiff *et al.*, 1979). Shiff and colleagues including

Clarke also demonstrated that mollusciding of irrigation canals using niclosamide drip-feed methods every 6-8 months as well as regular treatment of drains with the molluscicide also significantly reduced the risk of infection with *S. mansoni* and *S. haematobium* in children (Shiff *et al.*, 1973). They also conducted an economic costing of the work as well as operational assessment concluding that only 10% level of surveillance and incidence check of sentinel sites within the irrigation was sufficient to provide informative monitoring and evaluation of the efficacy and long-term effects of the molluscicide control efforts (Shiff *et al.*, 1973). In a recent meta-analysis of 35 molluscicide studies including several from Zimbabwe, King and colleagues reported that on average mollusciding reduced the odds of infection by 77% with the effects increased if mollusciding was integrated with antihelminthic treatment of the human population, while incidence was reduced by 64%, but interestingly antihelminthic treatment did not influence the incidence of infection (King *et al.*, 2015).

The irrigation mollusciding study in Zimbabwe in 1973 showed that controlling schistosomiasis via this method cost between USD54,800 and USD 55,500 for a 30,000 ha irrigation scheme (Shiff *et al.*, 1973). Thus, although the control efforts integrating antihelminthic treatment and mollusciding proved effective, not surprisingly, the vector control was not sustainable, either economically or environmentally. To overcome the toxic effects of niclosamide on plants, other snails (potential competitors) and fish, biological control strategies presented an attractive alternative. In Zimbabwe different biological interventions have been investigated; these included molluscicides derived from the plants *Phytolacca dodecandra* and *Jatropha curcas* with mixed efficacy and mixed community uptake (Madhina *et al.*,

1996). In addition snail predators have also been invoked in the form of ducks and fish (Chimbari, 2012). Poaching of the former (which were non-indigenous duck species) and low efficacy of the latter reduced the uptake of these interventions. Introduction of non-host competitor snails did not have a significant effect on the population of the intermediate host snails (Chimbari, 2012).

Engineering strategies to control schistosomiasis

Engineering and environmental interventions cannot be stressed enough in schistosome control. Long before the schistosome epidemic in Senegal in the 1990s, following the damming of the Senegal river to provide water for a sugar irrigation scheme in Richard Toll (Stelma *et al.*, 1993), Zimbabwe had experienced a schistosome epidemic as a result of the construction of the Kariba Dam in the late 1950s (Hira, 1970). The lessons learnt from this episode inspired the collaborative work between health professionals and engineers in the design of dam and irrigation schemes. A very successful example of this was the Mushandike Irrigation Scheme initiated in 1986 (Chimbari, 1991). Irrigation canals were lined to facilitate fast movement of irrigation water to dislodge snails and also comprised features which flushed out and trapped snails. Toilets were constructed along the scheme in a matrix ensuring that the workers were always nearer to a toilet than to the bush (Chimbari, 2012). This programme was so successful that despite the high costs of the design, the model was adopted by the Department of Irrigation in Zimbabwe as standard for all small irrigation schemes (Chimbari, 2012).

ZIMBWBWE'S NATIONAL SCHISTOSOMIASIS CONTROL PROGRAMME

447 With the history of commitment to schistosome research and control demonstrated
448 by the previous studies, it is not surprising that Zimbabwe has conducted regular
449 national surveys of schistosome infection in humans as well as the distribution and
450 infectivity of intermediate host snails. The first comprehensive national
451 schistosomiasis survey was conducted in 1982, followed by the second in 1992 (see
452 (Chimbari, 2012) for details). We conducted the third and most recent national
453 schistosomiasis and soil transmitted helminth (SHT) survey in Zimbabwe in 2010
454 (Midzi *et al.*, 2014). One of the changes that occurred in the intervening period
455 between the 1992 survey and our survey was the momentum from the global
456 initiative to control schistosomiasis following the 2001 World Health Assembly
457 Resolution 54.19 to treat at least 75% of all school aged children who are at risk of
458 morbidity from schistosomiasis and Soil transmitted helminths (STH) by the year
459 2010. This, coupled with the wider availability and significant reduction in price of the
460 antihelminthic drug PZQ, made for a suitable environment to implement a national
461 schistosomiasis control programme in Zimbabwe. When we published the findings of
462 our national schistosomiasis survey, we also included a national plan of action
463 incorporating the treatment strategies recommended by the WHO (Midzi *et al.*,
464 2014). Following the national helminth survey, we contributed to the formulation of
465 the national schistosomiasis and helminth control policy in Zimbabwe through a
466 workshop hosted by the Ministry of Health in Zimbabwe. To continue with the
467 research legacy of Zimbabwe and inter-sectorial collaborative approaches to
468 schistosome control, this policy document highlighted among others, the importance
469 of continued scientific research, dialogue between engineers and health
470 professionals, dialogue between the ministries of health and education both for
471 health education and implementation of the national control programmes and

community involvement. Following operational results from studies we conducted in schools in Zimbabwe and input from several stakeholders the Ministry of Health in Zimbabwe launched a the 5-year national schistosomiasis and soil transmitted control programme in September 2012. This programme is targeting just under 5 million primary school children throughout the country, treating all children annually regardless of the transmission level in the areas. The full evaluation of the programme will be conducted at the end of the 5 year programme but early indications from the sentinel monitoring and evaluation sites are that there has been a reduction in infection levels.

FUTURE PLANS AND CONCLUSION

The question for Zimbabwe and all other countries currently implementing national schistosomiasis control programmes is what happens after the 5 years of MDA? From our immuno-epidemiology and quantitative studies in Zimbabwe, we have predicted that cessation of MDA programmes may result in a rebound in infection to levels higher than pre-treatment levels (Mitchell *et al.*, 2014). Indeed earlier studies in Zimbabwe in the 1990s showed infection levels returning to pre-intervention levels when control (even strategies using integrated methods applied for 5-year periods were ceased) (Chimbari, 2012). These studies indicate that there is need for sustained control efforts and long-term planning to avoid areas of refugia for the parasites as well to facilitate the move from morbidity control to controlling transmission. There is also need for inclusive controls strategies if elimination is a realistic goal for schistosomiasis. Targeting primary school children while leaving the preschool children and adults will not lead to elimination of the diseases especially

where untreated infections in adulthood can become chronic with complications only addressed by surgery.

There are still areas needing more research including diagnostics, therapeutics and operational aspects as recently highlighted by Secor (Secor & Montgomery, 2015). There is also need for a more integrated approach to disease control involving dialogue between different sectors such as social scientists, engineers, architects (to enable building of human cities and dwellings that interrupt parasite transmission) and economists to come up with sustainable solutions to schistosome control. The current generation of schistosome researchers working in Zimbabwe aims to contribute to this knowledge base and strengthen the legacy of putting research before policy and implementation in schistosomiasis control.

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758 LIST OF FIGURES

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760 Fig 1: Photographs of VIP Latrines from Zimbabwe A) family setting, B) school
761 setting

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763 Fig 2: Photograph of a borehole using the bust pump design (village setting)

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For Peer Review